UNCLASSIFIED

AD 439310

DEFENSE DOCUMENTATION CENTER

FOR

SCIENTIFIC AND TECHNICAL INFORMATION

CAMERON STATION, ALEXANDRIA, VIRGINIA



UNCLASSIFIED

MOTICE: When government or other drawings, specifications or other data are used for any purpose other than in connection with a definitely related government procurement operation, the U.S. Government thereby incurs no responsibility, nor any obligation whatsoever; and the fact that the Government may have formulated, furnished, or in any way supplied the said drawings, specifications, or other data is not to be regarded by implication or otherwise as in any manner licensing the holder or any other person or corporation, or conveying any rights or permission to manufacture, use or sell any patented invention that may in any way be related thereto.

Quarterly REPORT OF INVESTIGATION

Quarterly REPORT OF INVESTIGATION

Quarterly REPORT OF INVESTIGATION

AD NO. PILE COPY

STUDIES ON THE MODE OF ACTION OF ANTIBACTERIAL DRUGS

RESPONSIBLE INVESTIGATOR

Dr. Katsuhiko Tago,

7/98800 Kitasato Instituto
Solya, (Japan)

U.S. Army Research & Development Group (9852) (Far East)

Office of the Chief of Research and Development
United States Army
APO 343

4 3 9 3 1 0

Quarterly No. 1 Report on Contract No. DA-92-557-FEG-35573 Inclusive Dates | Nov. 1961 to 31 Jan. 1962

Subject of Investigation

OF ANTIBACTERIAL ERUGS

Responsible Investigator

Dr. Katsuhiko Tego

Assist. Chief. Tobercologis Section Kitaseto Institute Tokyo, Japan Katsuhiko Tago, Yukiyoshi Yajima and Toru Hayakawa From Tuberculosis Section, Kitasato Institute Tokyo, Japan

A new antibiotic, mitomycin, has been isolated in our institute by Dr. Hata, from a new species of streptomycete, Streptomyces caespitosus. This substance was detected in the course of the screening program for the antitumor agents against the Phrlich carcinoma, antitumor activity was found to be in correlation with the antibacterial activity against B. subtilis PCI 219. The extraction of the active principle was carried out depending upon this antibacterial ctivity. result, the isolated principle was further separated into 3 active fractions. The names of mitomycin A. B and C were given to the respective fractions. The mitomycins showed strong inhibitory activities against a variety of gram-positive and gram-negative bacteria in vitro. as shown in Table 1. Among them, mitomycin A may be one of the most potential applibiotics which have so far been reported,

Table 1. Antibactorial Spectrum of Mitomycips

Minimum Inhibitory Concent				
Test Creanisms	4 जल्द	В под		
Staphylococcus aureus 209-p	0.005	0.1		
Diplococcus pneumoniae	0.01	0.1		
Streptococcus hemolyticus	0.01	0.1		
Corynebacterium diphtheriae	0.001 0.15	0.1 0.2		
Escherichia coli Klebsiella pneumoniae	0.15	0.005		
Salmonella typhi	0.5	0.8		
Shigella dysenteriae	0.15	6.8		
Bacterium subtilis ECI 219	0.001	0.025		

- 1. Inhibition of bacterial protein synthesis by mytomycin.
 - a. Materials and Methods.
 - The organism. The organism used for this work was Staphylococcus aureus strain 200-p grown for 18 hrs. at 37°C on normal agar media and the cells were harvested and washed once with 0.1 M phosphate buffer pH 6.25 and resuspended the same buffer. The cell suspensions were subjected to freezing and thawing twenty times. The disintegrated material was centrifuged at 1,000 r.p.m. for 6 min. This sediment was called Fraction I and the supernatant was resuspended 10 ml. of the buffer and recentrifuged at 6,000 r.p.m. for 30 min. and the sediment was refared to Fraction II, and the supernatant, Fraction III.
 - (2) Incubation mixtures. Incubation were carried out in 15 ml. centrifuge tubes containing the following: 0.1 ml. 0.06M Adenosin triphosphate; 0.4 ml. Hexose diphosphate solution containing the equivalent of 100 mg, barium salt/ml.; 0.5 ml. amino acid mixture A as disoribed the previous paper, 0.5 ml. the solution of antibacterial drugs and 0.5 ml. bacterial cell fraction.
 - Precipitation and estimation of proteinnitrogen. After incubating the tubes in a
 water bath at 3700 for 5 hrs, the same
 volume of 10% trichloracetic acid (AMO)
 was added to each tube and kept in ice box
 overnight. After centrifugation at 3,500
 r.p.m. for 20 min., the sediments were
 washed once with 5% MCA. The nitrogen
 content of this precipitate was determined
 by digestion in microhjeldahal apparatus
 and colorimetric after nesslerization.

b. Results

Table 2. Inhibition of protein synthesis by Mitomyoin

	Concentration mcg/ml	Inhibition
Mitomycin O	50 10 5	0
Chloramphericol	50 10 5	1.00 1.00 78
Leucomycin	50 10	97 94
	5 1	94 90

- (1) In Table 2 are represented the degree of inhibition produced by a variety of inhibitors tested on the synthesis of protoin. Inhibition is expressed as percentage rate of protein synthesis in the absence of added inhibitor. Mean value of controls without inhibitor is 60 mcg/mg dry wt. disrupted cell.
- (2) Mitomyoin C, although its high bacteriostatic activity, did not show any inhibitory power against the bacterial proteinsynthesis.
- (3) Chloramphenicol and Ieuconycin showed high inhibitory activity against bacterial protein synthesis. The degree of inhibition of protein-synthesis was well correlated with their respective antibacterial activity.

- 2. Inhibition of beta-galactosidase induction by mitomycin,
 - Material and Methods. a.
 - (1) The organism. The organism employed for this experiment was also Staphylococcus aureus 209-p and three fractions of the disrupted cells were prepared quite same as the experiment 1.
 - (2) The drugs. Mitomycin, Chloramphenicol and Leucomycin were tested. Concentration of the antibacterial agents were adjusted to obtain a suitable level (100-0.05 mcg per ml. of final) in reactions mixture. The solution of drugs were stored in a refrigerator during the experiment and the activity of the compounds did not show any change.
 - (3) The enzyme induction and assay of its activity. The ability of cellular suspensions to synthesize beta-galactosidase was determined with the O-hitrophenylbeta-d-galactoside method of Lederberg (1950) modified by Hurwitz et al. (1958). 0.2 ml, of each fractions contained in 15 ml. centrifuge tubes were added with l ml. of drug solution, 0.4 ml. of mixture B modified from medium A as follows: WazHPO4 3.2%; KH2704 0.8%; Na-oitrate 0.2%; MgSO4 0.04%; NH401 0.4%; Glucose 0.8% replaced with galaotose as an inducer. The tubes were incubated at 3700 for 5 hrs. in a water bath. At the end of the incubation a few drop of toluol were added to make cryptic enzyme accessible to substrate. Then one ml. of M/200 0-mitrophenyl-beta-d-galactoside (ONPG) solution and A ml. of 0.2 M phosphate buffer pH 7.5 were added to the incubation mixture to measure enzyme activity and the tubes were reheated to 3700 and kept at this temperature for 15 min.. This reaction was stopped

after a given time interval by addition of 1 ml. of 1 M sodium Carbonate. After the centrifugation of the tubes to clarify the reaction mixture, the intensity of yellow colour developed in cell free fluid resulting from liberation of o-nitrophenyl by cleavage of the ONFG was estimated Pulfrich photometer with No. 53 filter.

h. Result.

Table 3. Inhibition of induced beta-galactosidese synthesis by Mitomycin

Mitomycin	Leucomycin	Chloramphenicol	Per cent inhibition
mag/ml			%
5.0	-	-	100
1.0	-	-	100
0.5	-	-	60
0.1	-	••	11
0.05			0
	mog/ml		- of
-	6.3	•••	100
-	3.2	-	100
-	1.6		69.2
-	0.8	.~	38.4
-	0.4 0.2	L-1	5.1
	0.2		0
	0.05 0.05	-	0
برحضت مستسبي		100 g Zm?	
~	**	6.3	100
-	-	3.2	100
-	-	1.6	83.8
-		nog/ml 6.3 5.2 1.6 0.8	50.2
-	-	0.4	16.2
-	-	0.2	10.8
-	~	0.1	9.0
		0.05	0

- (1) The result obtained were indicated in the Table 2. In this experiment the effects of graded concentration of the antibacterial drugs on betagala-ctosidase induction of Staphylococus surevs were observed a range of 0.05 to 5.0 mcg/ml. Activity of beta-galactosidase is shown in percent and the value of 100 per cent means the normal formation of the enzyme that is obtained in the abscence of inhibitors. If the complete inhibition of the enzyme induction occurs, the value may reach to zero.
- (2) Induction of beta-galactosidase of Fraction II, coll-free extract, was completely inhibited by mitomycin at the concentration 1.0 mcg/ml.. Fifty percent inhibition dose was 0.45 mcg/ml..
- (3) Fifty percent inhibition dose of Chloramphenical and alpha-Bromocin-namaldehyd is 0.1 mag/ml. and 0.8 mag/ml. respectivly.